1 2	REGULATION 5.20 Methodology for Determining Benchmark Ambient Concentration Toxic Air Contaminant	of a						
3 4	Air Pollution Control District of Jefferson County Jefferson County, Kentucky							
5	Relates To: KRS Chapter 77 Air Pollution Control							
6	Pursuant To: KRS Chapter 77 Air Pollution Control							
7	Necessity and Function: KRS 77.180 authorizes the Air Pollution Control Board to adopt	and						
8	enforce all orders, rules, and regulations necessary or proper to accomplish the purposes of l							
9	Chapter 77. This regulation establishes the methodology for determining the benchmark amb	oient						
10	concentration for a toxic air contaminant.							
11	SECTION 1 Use of Benchmark Ambient Concentration							
12	A benchmark ambient concentration for a toxic air contaminant developed pursuant to this regula	ation						
13	shall be used in Regulation 5.21 Environmental Acceptability for Toxic Air Contaminan							
14	determine environmental acceptability.							
15	SECTION 2 Determination that a Toxic Air Contaminant is a Carcinogen							
16	2.1 A toxic air contaminant (TAC) shall be determined to be a carcinogen if any of the follow	ving						
17	provisions is met:							
18	2.1.1 A carcinogenic unit risk estimate, or alternatively, a concentration representative	of a						
19	specified level of additional lifetime cancer risk, for the TAC is included in any o	f the						
20	information sources identified in section 3.3,							
21	2.1.2 The TAC is listed as either "known to be a human carcinogen" or "reasonably anticip	ated						
22	to be a human carcinogen" in the most recent Report on Carcinogens published by	y the						
23	National Toxicology Program pursuant to Section 301(b)(4) of the Public Health Ser	vice						
24	Act as Amended by Section 262, PL 95-622, available on the Interne	t at						
25	"http://ehp.niehs.nih.gov.roc", or							
26	2.1.3 The District determines that the TAC should be considered to be a carcinogen bec	ause						
27	there is sufficient, credible information that any of the following criteria is met:							
28	2.1.3.1 Known to be a human carcinogen: There is sufficient evidence of carcinogen	icity						
29	from studies in humans which indicates a causal relationship between exposure t	o the						
30	agent, substance, or mixture and human cancer,							
31	2.1.3.2 Reasonably anticipated to be a human carcinogen:							
32	2.1.3.2.1 There is limited evidence of carcinogenicity from studies in humans, w	hich						
33	indicates that causal interpretation is credible, but that alternative explanati							
34	such as chance, bias, or confounding factors, could not adequately be exclu							
35	2.1.3.2.2 There is sufficient evidence of carcinogenicity from studies in experime							
36	animals which indicates there is an increased incidence of malignant	or a						
37	combination of malignant and benign tumors: (1) in multiple species of							
38	multiple tissue sites, or (2) by multiple routes of exposure, or (3) to an unu	sual						
39	degree with regard to incidence, site, or type of tumor, or age at onset, or							
40	2.1.3.2.3 There is less than sufficient evidence of carcinogenicity in humans or laborated the sufficient evidence of carcinogenicity in humans or laborated the sufficient evidence of carcinogenicity in humans or laborated the sufficient evidence of carcinogenicity in humans or laborated the sufficient evidence of carcinogenicity in humans or laborated the sufficient evidence of carcinogenicity in humans or laborated the sufficient evidence of carcinogenicity in humans or laborated the sufficient evidence of carcinogenicity in humans or laborated the sufficient evidence of carcinogenicity in humans or laborated the sufficient evidence of carcinogenicity in humans or laborated the sufficient evidence of carcinogenicity in humans or laborated the sufficient evidence of carcinogenicity in humans or laborated the sufficient evidence of carcinogenicity in humans or laborated the sufficient evidence of carcinogenicity in humans or laborated the sufficient evidence of carcinogenicity in humans or laborated the sufficient evidence of carcinogenic evidence evidence of carcinogenic evidence eviden							
41	animals, however; the agent, substance, or mixture belongs to a well defi	ned,						

[If adopted, this would be a new regulation]

structurally-related class of substances whose members are listed in the most recent *Report on Carcinogens* published by the National Toxicology Program as either a known to be human carcinogen or reasonably anticipated to be human carcinogen, or there is convincing relevant information that the agent acts through mechanisms indicating it would likely cause cancer in humans.

- 2.2 In making a determination pursuant to section 2.1.3, the following provisions shall apply:
- 2.2.1 Conclusions regarding carcinogenicity in humans or experimental animals are based on scientific judgment, with consideration given to all relevant information. Relevant information includes, but is not limited to, dose response, route of exposure, chemical structure, metabolism, pharmacokinetics, sensitive sub populations, genetic effects, and other data relating to mechanism of action or factors that may be unique to a given substance. This applies to both the "known to be a human carcinogen" and the "reasonably anticipated to be a human carcinogen" categories, and
- 2.2.2 For an agent to be determined "known to be a human carcinogen," evidence from studies of humans is required. This may include traditional cancer epidemiology studies, data from clinical studies, or data derived from the study of tissues from humans exposed to the substance in question and useful for evaluating whether a relevant cancer mechanism is operating in humans.

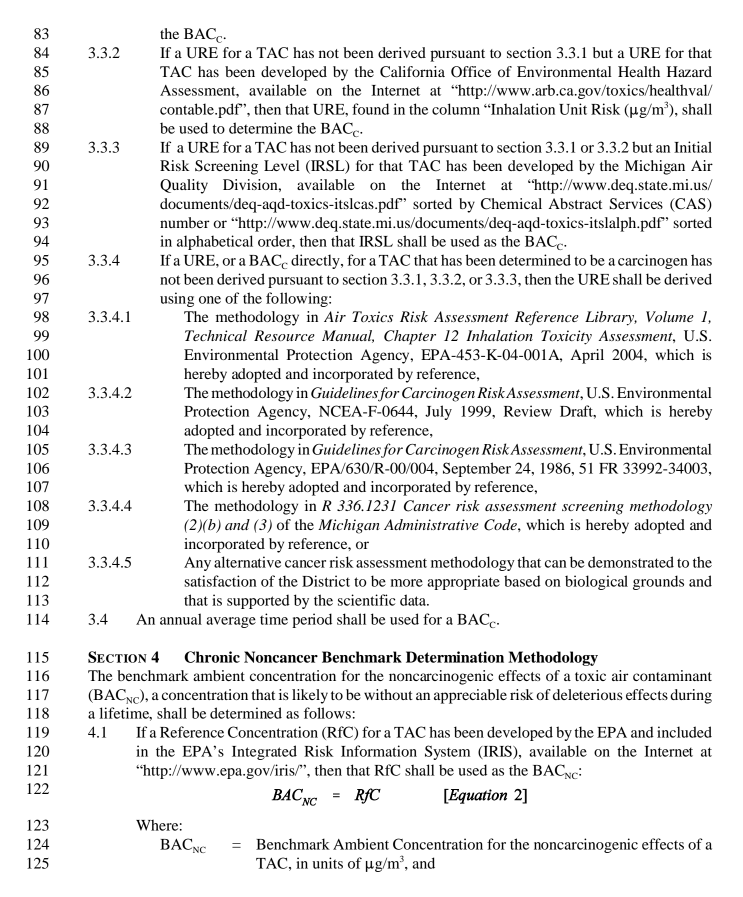
SECTION 3 Cancer Risk Benchmark Determination Methodology

3.1 The benchmark ambient concentration for a toxic air contaminant (TAC) determined to be a carcinogen (BAC_C) shall be calculated as follows:

$$BAC_C = \frac{1 \otimes 10^{-6}}{URE}$$
 [Equation 1]

Where:

- BAC_C = Benchmark Ambient Concentration for a carcinogen, a concentration representative of an additional lifetime cancer risk of 1 in 1,000,000 ($1 \otimes 10^{-6}$), in units of micrograms per cubic meter (μ g/m³),
- URE = Unit Risk Estimate Additional lifetime cancer risk occurring in a population in which all individuals are exposed continuously for life (70 years) to a concentration of $1 \mu g/m^3$ of the chemical in the air they breathe, in units of $(\mu g/m^3)^{-1}$. The URE shall be determined according to the methodology in section 3.3, and
- $1 \otimes 10^{-6}$ = An upper bound additional lifetime cancer risk of 1 in 1,000,000.
- 3.2 Alternatively, if in any of the sources of information identified in section 3.3, the concentration of a carcinogen, expressed in $\mu g/m^3$, that is representative of an additional lifetime cancer risk of $1 \otimes 10^{-6}$ is identified instead of the URE, then the BAC_C is that identified concentration. The URE can be calculated by using Equation 1.
- 3.3 The following provisions shall apply to the derivation of a unit risk estimate (URE), or alternatively a BAC $_{C}$ directly, for a TAC determined to be a carcinogen:
- 3.3.1 If a URE for a TAC has been developed by the U.S. Environmental Protection Agency (EPA) and included in the EPA's Integrated Risk Information System (IRIS), available on the Internet at "http://www.epa.gov/iris/", then that URE shall be used to determine



126 RfC = Reference Concentration, in units of $\mu g/m^3$.

127 A 24-hour average time period shall be used for a BAC $_{NC}$ determined pursuant to section 4.1.

4.2 If a BAC_{NC} for a TAC has not been determined pursuant to section 4.1 but a Reference Exposure Level (REL) for that TAC has been developed by the California Office of Environmental Health Hazard Assessment, available on the Internet at "http://www.arb.ca.gov/toxics/healthval/contable.pdf", then that REL, found in the column "Chronic Inhalation ($\mu g/m^3$), shall be used as the BAC_{NC}:

 $BAC_{NC} = REL$ [Equation 3]

134 Where:

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= Benchmark Ambient Concentration for the noncarcinogenic effects of a 135

TAC, in units of $\mu g/m^3$, and

= Reference Exposure Level, in units of $\mu g/m^3$. 137 **REL**

138 A 24-hour average time period shall be used for a BAC $_{NC}$ determined pursuant to section 4.2.

4.3 139 If a BAC_{NC} for a TAC has not been determined pursuant to section 4.1 or 4.2 but an Oral 140 Reference Dose (RfD) for that TAC has been developed by the EPA and included in the EPA's IRIS, available on the Internet at "http://www.epa.gov/iris/", and data are not 141 142 available to indicate that oral-route to inhalation-route extrapolation is inappropriate, then 143 that RfD shall be used to calculate the BAC $_{NC}$ as follows:

 $BAC_{NC} = Oral \ RfD \otimes \frac{70 \ kg}{20 \ \frac{m^3}{dgy}}$ [Equation 4]

Where: 145

= Benchmark Ambient Concentration for the noncarcinogenic effects of a

147 TAC, in units of $\mu g/m^3$,

= Reference Exposure Level, in units of µg/kg-day, RfD 148

149 70 kg = The average body weight of a human, and

150 $20 \text{ m}^{3}/\text{day} =$ The average daily inhalation rate for a human.

151 A 24-hour average time period shall be used for a BAC $_{NC}$ determined pursuant to section 4.3.

4.4 If a BAC_{NC} for a TAC has not been determined pursuant to section 4.1 to 4.3 but an Initial Threshold Screening Level (ITSL) for that TAC has been developed by the Michigan Air Quality Division, available on the Internet at "http://www.deq.state.mi.us/ documents/deg-agd-toxics-itslcas.pdf" sorted by Chemical Abstract Services (CAS) number or "http://www.deq.state.mi.us/documents/deq-aqd-toxics-itslalph.pdf" sorted in alphabetical order, then that ITSL shall be used as the BAC $_{NC}$:

> $BAC_{NC} = ITSL$ [Equation 5]

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[If adopted, this would be a new regulation]

159 Where: = Benchmark Ambient Concentration for the noncarcinogenic effects of a 160 BAC_{NC} 161 TAC, in units of $\mu g/m^3$, and = Initial Threshold Screening Level, in units of $\mu g/m^3$. 162 ITSL 163 The average time period as listed for a specific ITSL shall be used for a BAC_{NC} determined pursuant to section 4.4. 164 If a BAC_{NC} for a TAC has not been determined pursuant to section 4.1 to 4.4 but an 165 4.5 occupational exposure level (OEL) exists for that TAC, then the OEL may be used to 166 167 calculate the BAC_{NC} as follows: 168 $BAC_{NC} = \frac{OEL}{100}$ [Equation 6] 169 Where: = Benchmark Ambient Concentration for the noncarcinogenic effects of a 170 BAC_{NC} TAC, in units of $\mu g/m^3$, 171 = Occupational Exposure Level, that, for the TAC, is the lowest value of 172 OEL 173 either the National Institute of Occupational Safety and Health (NIOSH)-174 recommended exposure level listed in current edition of the NIOSH 175 pocket guide to chemical hazards or the time-weighted average or ceiling Threshold Limit Value (TLV) listed in the current edition of the 176 177 American Conference of Governmental and Industrial Hygienists 178 Threshold Limit Value (TLV) booklet, in units of $\mu g/m^3$, and = A composite safety factor to account for differences in susceptibility 179 100 180 between the healthy, adult worker population compared to the general population that is more diverse and may contain individuals or 181 182 subpopulations more sensitive to the effects of the toxic air pollutant (safety factor of 10). Additionally, the composite safety factor accounts 183 for the difference in exposure duration (in hours per week and years 184 working versus a lifetime) for the worker population compared to the 185 186 general population: 187 $\frac{1}{10} \otimes \frac{40 \text{ hours/week}}{168 \text{ hours/week}} \otimes \frac{30 \text{ years}}{70 \text{ years}} \approx \frac{1}{100}.$

An 8-hour average time period shall be used for a BAC $_{\rm NC}$ determined pursuant to section 4.5 based upon a time-weighted OEL and a 1-hour average time period shall be used for a BAC $_{\rm NC}$ determined pursuant to section 4.5 based upon a ceiling OEL.

191 4.6 If a BAC_{NC} for a TAC has not been determined pursuant to section 4.1 to 4.5 but a 7-day, 192 inhalation, no observed adverse effect level (NOAEL) or lowest observable adverse effect 193 level (LOAEL) is available for that TAC, then the NOAEL or LOAEL may be used to 194 calculate the BAC_{NC} as follows:

$$BAC_{NC} = \frac{NOAEL}{35 \otimes 100} \otimes \frac{Hr \ Exposed / \ Day}{24 \ Hr / \ Day} \qquad [Equation 7]$$

$$BAC_{NC} = \frac{LOAEL}{35 \otimes 100 \otimes UF} \otimes \frac{Hr \ Exposed / \ Day}{24 \ Hr / \ Day} \qquad [Equation 8]$$

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$$Equation 10$$

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[If adopted, this would be a new regulation]

227		35	=	A safety factor to account for using a NOAEL or LOAEL from a 7-day
228				exposure period to estimate a NOAEL or LOAEL for a lifetime study,
229		100	=	A standard composite safety factor comprised of a safety factor of 10 to
230				account for differences between animals and humans and a safety factor
231				of 10 to account for the differences between individuals in the human
232				population,
233		UF	=	Uncertainty Factor, a value from 1 to 10, applicable when using a
234				LOAEL (lowest effect) instead of a NOAEL (no effect), determined by
235				the District on a case-by-case basis, considering the type and severity of
236				effect. For example, a value of 1 would be used when the lowest effect
237				was a skin rash; a value of 10 would be used when the lowest effect was
238				death,
239		\mathbf{W}_{A}	=	Body weight of experimental animal in kilograms (kg),
240		I_A	=	Daily inhalation rate of experimental animal in m³/day,
241		b	_	Absorption efficiency (percent absorbed) by the oral route of exposure,
242		Ü	_	and
243		a	=	Absorption efficiency (percent absorbed) by the inhalation route of
244		u	_	exposure.
277				exposure.
245		If approved b	v the	e District, the BAC _{NC} may be determined on a case-by-case basis using an
246				ral LOAEL from repeated dose studies other than 7-day studies.
210		Oldi I (Oldi	01 0	Tai Destille from repeated dose stadies other than 7 day stadies.
247		An annual av	erage	e time period shall be used for a BAC_{NC} determined pursuant to section 4.7.
248	4.8	If a BAC _{NC}	for a	a TAC has not been determined pursuant to section 4.1 to 4.7 but an
249				rom a study that is 4 or more hours in duration is available for that TAC,
250				y be used to calculate the BAC_{NC} as follows:
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			1	$BAC_{NC} = \frac{LC_{50}}{500 \otimes 100}$ [Equation 11].
252		Where:		
253		BAC_{NC}	=	Benchmark Ambient Concentration for the noncarcinogenic effects of a
254		INC		TAC, in units of $\mu g/m^3$,
255		LC_{50}	=	Concentration of material used in an inhalation study that causes death of
256		50		50% of the group of test animals when administered as a single dose in
257				a specific time period, in units of $\mu g/m^3$,
258		500	=	A factor to account for using an LC_{50} to estimate a no observed adverse
259		500	_	effect level (NOAEL) for a lifetime study, and
200		100		A to 1 1 1 Constitution of the Constitution of

An annual average time period shall be used for a BAC_{NC} determined pursuant to section 4.8.

= A standard composite safety factor comprised of a safety factor of 10 to

account for differences between animals and humans and a safety factor of 10 to account for the differences between individuals in the human

population.

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265 4.9 If a BAC $_{NC}$ for a TAC has not been determined pursuant to section 4.1 to 4.8 but an LC $_{50}$ from a 1-hour inhalation study is available for that TAC, then the 1-hour LC $_{50}$ may be used to calculate the BAC $_{NC}$ as follows:

 $BAC_{NC} = \frac{(1-Hr) LC_{50}}{500 \otimes 100 \otimes 40}$ [Equation 12].

269	Where:		
270	BAC_{NC}	=	Benchmark Ambient Concentration for the noncarcinogenic effects of a
271			TAC, in units of $\mu g/m^3$,
272	LC_{50}	=	Concentration of material used in an inhalation study that causes death of
273			50% of the group of test animals when administered as a single dose in
274			a specific time period, in units of $\mu g/m^3$,
275	500	=	A factor to account for using an LC ₅₀ to estimate a no observed adverse
276			effect level (NOAEL) for a lifetime study,
277	100	=	A standard composite safety factor comprised of a safety factor of 10 to
278			account for differences between animals and humans and a safety factor
279			of 10 to account for the differences between individuals in the human
280			population, and
281	40	=	A safety factor to account for the uncertainty of using a one-hour
282			inhalation LC_{50} compared to an exposure duration of four hours or more.

An annual average time period shall be used for a BAC $_{NC}$ determined pursuant to section 4.9.

4.10 If a BAC_{NC} for a TAC has not been determined pursuant to section 4.1 to 4.9 but an animal oral LD₅₀ is available for that TAC, then the LD₅₀ may be used to calculate the BAC_{NC} as follows:

$$BAC_{NC} = \frac{LD_{50} (mg/kg)}{500 \otimes 100 \otimes 40 \otimes 0.167} \otimes \frac{W_A}{I_A}$$
 [Equation 13].

288	Where:		
289	BAC_{NC}	=	Benchmark Ambient Concentration for the noncarcinogenic effects of a
290			TAC, in units of $\mu g/m^3$,
291	LD_{50}	=	Amount of material administered in a single dose by a route other than
292			inhalation, e.g., oral, that causes death of 50% of the group of test
293			animals, in units of μg/kg,
294	500	=	A factor to account for using an LC_{50} to estimate a no observed adverse
295			effect level (NOAEL) for a lifetime study,
296	100	=	A standard composite safety factor comprised of a safety factor of 10 to
297			account for differences between animals and humans and a safety factor
298			of 10 to account for the differences between individuals in the human
299			population,
300	40	=	A safety factor to account for the uncertainty of estimating an LC ₅₀ from
301			an LD_{50} ,

302 303 304	0.167 = A factor to convert the daily dose to a 4-hour time frame $(4 \div 24 = 0.167)$, W_A = Body weight of experimental animal in kilograms (kg), and I_A = Daily inhalation rate of experimental animal in m³/day.
305 306	An annual average time period shall be used for a BAC_{NC} determined pursuant to section 4.10.
307 308	4.11 If a BAC $_{\rm NC}$ for a TAC has not been determined pursuant to section 4.1 to 4.10, then the BAC $_{\rm NC}$ shall be the default value:
309	$BAC_{NC} = 0.04 \mu g/m^3$ [Equation 14].
310 311 312	Where: $BAC_{NC} = Benchmark \ Ambient \ Concentration \ for \ the \ noncarcinogenic \ effects \ of \ a \\ TAC, \ in \ units \ of \ \mu g/m^3.$
313 314	An annual average time period shall be used for a $BAC_{\text{\tiny NC}}$ determined pursuant to section 4.11.
315 316 317 318	Section 5 Consideration of Acute Noncancer Effects If the District believes that compliance with the BAC_{NC} over the applicable averaging time specified in Section 4 does not provide adequate protection from the acute effects of a TAC, then the District may establish a different BAC_{NC} and shorter averaging time that would provide adequate protection.
319 320 321 322 323	SECTION 6 Available Documents The District will maintain on its web page links to the documents identified as available on the Internet and maintain at its office a copy of all documents identified in this regulation. In addition, the District will maintain a current list of the benchmark ambient concentrations that have been developed pursuant to this regulation and maintain this current list on its web page.
324	Adopted v1/; effective